

FORM PTO-1390 (Modified)
(REV 11-98)

U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE

ATTORNEY'S DOCKET NUMBER

TRANSMITTAL LETTER TO THE UNITED STATES

203852US0PCT

DESIGNATED/ELECTED OFFICE (DO/EO/US)

U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR

CONCERNING A FILING UNDER 35 U.S.C. 371

09/786113

INTERNATIONAL APPLICATION NO.

PCT/FR99/02068

INTERNATIONAL FILING DATE

30 August 1999

PRIORITY DATE CLAIMED

01 September 1998

TITLE OF INVENTION

METHOD OF IMPROVING THE MECHANICAL STRENGTH OF AN INSULATION PRODUCT BASED ON MINERAL WOOL, INSULATION PRODUCT AND SIZING COMPOSITION

APPLICANT(S) FOR DO/EO/US

Jacky JOACHIM, et al.

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

1. ☒ This is a **FIRST** submission of items concerning a filing under 35 U.S.C. 371.
2. ☐ This is a **SECOND** or **SUBSEQUENT** submission of items concerning a filing under 35 U.S.C. 371.
3. ☒ This is an express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1).
4. ☒ A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.
5. ☒ A copy of the International Application as filed (35 U.S.C. 371 (c) (2))
 - a. ☐ is transmitted herewith (required only if not transmitted by the International Bureau).
 - b. ☒ has been transmitted by the International Bureau.
 - c. ☐ is not required, as the application was filed in the United States Receiving Office (RO/US).
- ☒ A translation of the International Application into English (35 U.S.C. 371(c)(2)).
- ☒ A copy of the International Search Report (PCT/ISA/210).
- ☒ Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371 (c)(3))
 - a. ☐ are transmitted herewith (required only if not transmitted by the International Bureau).
 - b. ☐ have been transmitted by the International Bureau.
 - c. ☐ have not been made; however, the time limit for making such amendments has NOT expired.
 - d. ☒ have not been made and will not be made.
- ☐ A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
- ☐ An oath or declaration of the inventor(s) (35 U.S.C. 371 (c)(4)).
- ☐ A copy of the International Preliminary Examination Report (PCT/IPEA/409).
- ☐ A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371 (c)(5)).

Items 13 to 20 below concern document(s) or information included:

13. ☐ An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
14. ☐ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
15. ☒ A **FIRST** preliminary amendment.
16. ☐ A **SECOND** or **SUBSEQUENT** preliminary amendment.
17. ☐ A substitute specification.
18. ☐ A change of power of attorney and/or address letter.
19. ☐ Certificate of Mailing by Express Mail
20. ☒ Other items or information:

Request for Consideration of Documents Cited in International Search Report

Notice of Priority

PCT/IB/304

PCT/IB/308

U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR 09/786113)		INTERNATIONAL APPLICATION NO. PCT/FR99/02068		ATTORNEY'S DOCKET NUMBER 203852US0PCT	
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21. The following fees are submitted:

BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)) :			CALCULATIONS PTO USE ONLY	
<input type="checkbox"/> Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO	\$1,000.00			
<input checked="" type="checkbox"/> International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO	\$860.00			
<input type="checkbox"/> International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO	\$710.00			
<input type="checkbox"/> International preliminary examination fee paid to USPTO (37 CFR 1.482) but all claims did not satisfy provisions of PCT Article 33(1)-(4)	\$690.00			
<input type="checkbox"/> International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(1)-(4)	\$100.00			
ENTER APPROPRIATE BASIC FEE AMOUNT =		\$860.00		
Surcharge of \$130.00 for furnishing the oath or declaration later than <input type="checkbox"/> 20 <input checked="" type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492 (e)).		\$130.00		
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE	
Total claims	21 - 20 =	1	x \$18.00	\$18.00
Independent claims	4 - 3 =	1	x \$80.00	\$80.00
Multiple Dependent Claims (check if applicable).			<input type="checkbox"/>	\$0.00
TOTAL OF ABOVE CALCULATIONS =				\$1,088.00
Reduction of 1/2 for filing by small entity, if applicable. Verified Small Entity Statement must also be filed (Note 37 CFR 1.9, 1.27, 1.28) (check if applicable).			<input type="checkbox"/>	\$0.00
SUBTOTAL =				\$1,088.00
Processing fee of \$130.00 for furnishing the English translation later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492 (f)).			+	\$0.00
TOTAL NATIONAL FEE =				\$1,088.00
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31) (check if applicable).			<input type="checkbox"/>	\$0.00
TOTAL FEES ENCLOSED =				\$1,088.00
			Amount to be:	\$
			refunded	\$
			charged	\$

☒ A check in the amount of **\$1,088.00** to cover the above fees is enclosed.

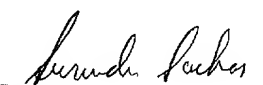
☐ Please charge my Deposit Account No. _____ in the amount of _____ to cover the above fees.
A duplicate copy of this sheet is enclosed.

☒ The Commissioner is hereby authorized to charge any fees which may be required, or credit any overpayment to Deposit Account No. **15-0030** A duplicate copy of this sheet is enclosed.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

SEND ALL CORRESPONDENCE TO:

Surinder Sachar
Registration No. 34,423



SIGNATURE

Norman F. Oblon

NAME

24,618

REGISTRATION NUMBER

March 1 2001

DATE

09/786113

203852US0PCT 01 MAR 2001

203852US0PCT

IN THE UNITED STATES PATENT & TRADEMARK OFFICE

IN RE APPLICATION OF: :

JACKY JOACHIM ET AL. :

SERIAL NO: NEW US PCT APPLN. : ATTN: APPLICATION BRANCH
(Based on PCT/FR99/02068)

FILED: HEREWITH :

FOR: METHOD OF IMPROVING THE
MECHANICAL STRENGTH OF AN
INSULATION PRODUCT BASED
ON MINERAL WOOL, INSULATION
PRODUCT AND SIZING COMPOSITION

PRELIMINARY AMENDMENT

ASSISTANT COMMISSIONER FOR PATENTS
WASHINGTON, D.C. 20231

SIR:

Prior to examination on the merits, please amend the above-identified application as follows.

IN THE SPECIFICATION

Please amend the specification as follows:

Page 1, before line 1, delete the title of the invention in its entirety, and insert therefor:

--METHOD FOR MAKING A FIBROUS INSULATING PRODUCT, SIZING
STUFF AND COMPOSITION--

IN THE CLAIMS

Please amend the claims as follows:

--3. (Amended) Method according to Claim 1, in which the latex is based on an aqueous dispersion or emulsion of a polymer carrying hydrophilic functional groups, especially hydroxyl, carboxyl or ester functional groups.

5. (Amended) Method according to Claim 3, in which the latex contains a polymer or copolymer which is of the vinyl type, especially a vinyl acetate homopolymer or copolymer, or of the acrylic type and/or which is derived from a carboxylic acid.

7. (Amended) Method according to Claim 1, in which the latex is based on an aqueous dispersion or emulsion of particles consisting of a polymer surrounded by a surfactant or by a protective colloid having hydrophilic functional groups, especially one based on polyvinyl alcohol or on cellulose.

9. (Amended) Method according to Claim 3, in which a water-repellent agent, such as a silicone or a fluorinated compound, is added to the latex.

10. (Amended) Method according to Claim 1, in which the latex is based on a polymer having a glass transition temperature T_g of less than 80°C and especially of less than 50°C .

11. (Amended) Method according to Claim 1, in which the latex is based on a polymer having a glass transition temperature T_g of greater than -5°C and especially of greater than 0°C .

12. (Amended) Method according to Claim 1, in which the solids content of the latex introduced is less than 5%, especially about 0.01 to 5%, by weight with respect to the weight of mineral wool.

13. (Amended) Method according to Claim 1, in which the latex is mixed with the size before application to the mineral wool.

14. (Amended) Method according to Claim 1, in which the latex is applied to the

mineral wool separately from the size.

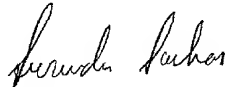
18. (Amended) Insulation product according to Claim 15, which has a density of at least 30 kg/m³, especially at least 50 kg/m³ and particularly at least 80 kg/m³.--

REMARKS

Claims 1-21 are active in the present application. The claims are amended to remove multiple dependencies. No new matter is added. An action on the merits and allowance of the claims is solicited.

Respectfully submitted,

OBLON, SPIVAK, McCLELLAND,
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Marked-Up Copy Serial No: _____ Amendment Filed on: _____
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--3. (Amended) Method according to Claim 1 [or 2], in which the latex is based on an aqueous dispersion or emulsion of a polymer carrying hydrophilic functional groups, especially hydroxyl, carboxyl or ester functional groups.

5. (Amended) Method according to Claim 3 [or 4], in which the latex contains a polymer or copolymer which is of the vinyl type, especially a vinyl acetate homopolymer or copolymer, or of the acrylic type and/or which is derived from a carboxylic acid.

7. (Amended) Method according to [one of Claims 1 to 3] Claim 1, in which the latex is based on an aqueous dispersion or emulsion of particles consisting of a polymer surrounded by a surfactant or by a protective colloid having hydrophilic functional groups, especially one based on polyvinyl alcohol or on cellulose.

9. (Amended) Method according to [any one of Claims 3 to 8] Claim 3, in which a water-repellent agent, such as a silicone or a fluorinated compound, is added to the latex.

10. (Amended) Method according to [any one of the preceding claims] Claim 1, in which the latex is based on a polymer having a glass transition temperature T_g of less than 80°C and especially of less than 50°C.

11. (Amended) Method according to [any one of the preceding claims] Claim 1, in which the latex is based on a polymer having a glass transition temperature T_g of greater than -5°C and especially of greater than 0°C.

12. (Amended) Method according to [any one of the preceding claims] Claim 1, in

which the solids content of the latex introduced is less than 5%, especially about 0.01 to 5%, by weight with respect to the weight of mineral wool.

13. (Amended) Method according to [any one of the preceding claims] Claim 1, in which the latex is mixed with the size before application to the mineral wool.

14. (Amended) Method according to [any one of Claims 1 to 12] Claim 1, in which the latex is applied to the mineral wool separately from the size.

18. (Amended) Insulation product according to [one of Claims 15 to 17] Claim 15, which has a density of at least 30 kg/m^3 , especially at least 50 kg/m^3 and particularly at least 80 kg/m^3 .--

METHOD OF IMPROVING THE MECHANICAL STRENGTH OF AN
INSULATION PRODUCT BASED ON MINERAL WOOL, INSULATION
PRODUCT AND SIZING COMPOSITION

5 The present invention relates to the techniques
used to manufacture insulation products, especially
thermal and/or acoustic insulation products, based on
mineral wool. It relates more particularly to an
10 improved size for the wool of which such a product is
composed, so as to improve the mechanical strength
after ageing of this product, particularly in a wet
medium.

These products, which may be based on glass or
rock wool, are usually in the form of rolled-up felts,
15 of relatively rigid panels, of shells or of blankets.

The manufacture of these insulation products
comprises the following steps:

- the glass or rock mineral composition is
melted in a suitable furnace;
- 20 - the molten mineral material is taken to a
fiberizing apparatus;
- the molten material is transformed into
filaments, especially using the known technique of
centrifugal drawing over rotors or of drawing through
25 perforated dishes, the filaments being generally drawn
under the action of a blast of gas;
- a sizing composition containing a
thermosetting resin is sprayed onto the wool thus
formed;
- 30 - the sized wool is gathered on a device for
taking it up in the form of a web; and
- the web is subjected to a heat treatment for
the purpose of curing the resin and the web is made
into the desired shape.

35 The properties required of the end-product
depend on each particular application, but it is
generally desired to obtain, in addition to the
insulating properties of the product, a number of
mechanical characteristics, such as dimensional

stability, puncture strength, tear strength, tensile strength and compressive strength.

Although it is important for these properties to remain good from the manufacture of the product until its use, in order to ensure the proper conditions for installing it on the site, it is also desirable for these properties to remain good for a certain length of time after the product has been put into place, in order to guarantee a long enough lifetime of the product.

Unfortunately, it sometimes happens that these products suffer a loss of mechanical properties after ageing, in particular when they are exposed to moisture, especially under conditions of high atmospheric condensation, for example in the case of products which are used in roofs and which are subjected to large thermal cycles.

It is also necessary, for example, to take care to avoid such losses when the composition of the mineral wool is particularly sensitive to water, for example when it is a mineral wool capable of dissolving in a physiological medium,

Document WO-A-97/21636 discloses how to improve the ageing strength in the presence of atmospheric moisture of artificial mineral fibres of the type soluble in a physiological medium (solubility of at least 20 nm/day at 7.5 pH at 37°C) by forming a coating on the fibres which comprises an ammonium hydrogenphosphate salt or a quaternary ammonium salt or an alkali metal salt, preferably diammonium hydrogenphosphate or ammonium dihydrogenphosphate.

The ageing strength in a wet medium is estimated in this document by deducing, from pH measurements, the degree of dissolution of the fibres in immersion tests simulating accelerated ageing under normal conditions of use. Optionally, in addition to the pH measurement, the erosion of the fibres is examined under a microscope.

However, this treatment proves to be insufficient for maintaining the mechanical properties of the product after exposure to the wet medium: by carrying out mechanical strength tests on these products based on mineral wool after accelerated ageing in a wet medium, the present inventors have observed that standard ammonium-based cationic agents were unable to limit the loss of mechanical properties after exposure to the wet medium and that a phosphate such as diammonium hydrogenphosphate even had a negative effect on the mechanical properties after ageing in a wet medium in the sense that the losses in properties were greater.

The object of the invention is to obviate these drawbacks and to improve the mechanical strength after ageing, particularly in a wet medium, of insulation products based on mineral wool, or in other words to reduce the loss of mechanical properties of these products after ageing, particularly in a wet medium.

This object, as well as others which will appear later, was achieved according to the invention by adding a latex to the size during the manufacture of the products.

For this purpose, the subject of the invention is a method of improving the mechanical strength after ageing, particularly in a wet medium, of an insulation product, especially a thermal and/or acoustic insulation product, based on mineral wool provided with a size comprising a thermosetting resin, especially a phenolic resin, in which method a latex is added to the size during the manufacture of the product.

Completely surprisingly, it has been found that although, in many cases, the addition of a latex to the size did not modify, or modified only slightly, the mechanical properties, or even degraded these properties right after manufacture, it was possible to achieve a remarkable reduction in the loss of properties after ageing, particularly in a wet medium, compared with a similar product not containing the

latex (standard product) and to achieve a final level of performance after ageing which is superior to the standard product.

Advantageously, this effect is observed on products based on mineral wool capable of being dissolved in a physiological medium (so-called "biosoluble" wool) which are products that ordinarily are quite sensitive to moisture, because of the quite high content of alkali metal oxides of the mineral material, often combined with a high content of boron oxide.

In the present application, the term "latex" should be understood to mean, in the usual manner, an aqueous dispersion or emulsion of one or more natural or synthetic, generally thermoplastic, polymer substances. The polymer or polymers may be self-emulsifiable, or else, if this is not so, the emulsion or dispersion is stabilized by suitable surfactants.

What proved to be advantageous, from the standpoint of the mechanical strength after exposure to the wet medium, are latices based on an emulsion or dispersion of a polymer phase carrying hydrophilic functional groups forming the interface with the aqueous phase. These functional groups are especially hydroxyl -OH, carboxyl -COOH or ester -COOR functional groups, where R denotes an alkyl group which may have especially from 1 to 5 carbon atoms. The ester functional groups are particularly preferred, especially the acetate functional group.

This result is completely surprising since it might be thought that a hydrophilic latex, by increasing the amount of water picked up by the product, would accelerate the loss of properties due to the wet medium, particularly in the case of products based on a so-called biosoluble mineral wool.

Without wishing to be bound by any scientific theory, it is possible that the hydrophilic character of the dispersed polymer phase of the latex gives the latter an advantageous affinity towards the mineral

material forming the wool, possibly because of the formation of polar bonds, making the latex act, as it were, as an adhesion primer for the resin. This is because it was found, in comparative tests of the tear
5 strength of the resin, that the prior application of a hydrophilic latex to the mineral wool gives greater adhesion of the resin of the size to the surface of the mineral material.

Whatever the situation, applying a hydrophilic
10 latex with the size during the manufacture of the insulation product results in products which withstand the ageing in a wet medium with smaller losses in mechanical properties.

In a preferred variant, the polymer itself
15 carries hydrophilic functional groups. Polymers in which each monomer carries at least one hydrophilic functional group prove to be advantageous in this regard, whether they be homopolymers derived from a single monomer or copolymers derived from at least two
20 different monomers. Nevertheless, the presence of a minor comonomer not carrying a hydrophilic functional group can be tolerated.

Advantageously, the latex contains a polymer or copolymer which is of the vinyl type or of the acrylic
25 type and/or which is derived from a carboxylic acid.

Most particularly preferred are latices of the vinyl type, particularly those having pendant ester functional groups, especially those based on vinyl acetate. Most especially preferred are latices based on
30 a polyvinyl acetate homopolymer, but mention may also be made, as advantageous latices, of those based on a copolymer of vinyl acetate and, especially, of a (meth)acrylic ester and/or acid, of a maleic ester, of an olefin and/or of vinyl chloride.

35 Other useful latices may be chosen from those containing an acrylic-type polymer, especially a silanized acrylonitrile/acrylic ester or styrene/acrylic ester or acid copolymer ("silanized" means copolymerized with a monomer having an

ethylenically unsaturated group carrying at least one silane or silanol functional group).

In certain cases, especially when the content of hydrophilic functional groups of the polymer is relatively low, the polymer may not have by itself a sufficiently hydrophilic character to be stable in emulsion and/or have a suitable affinity for the glass. Thus, the latex is advantageously such that the dispersed phase consists of a polymer surrounded by a protective colloid having hydrophilic functional groups (the whole forming a dispersed microparticle or nanoparticle), this colloid providing the desired hydrophilic functional groups on the surface of the suspended particle, i.e. at the interface with the aqueous phase.

In general, the colloid consists of one or more macromolecules; advantageously, it may be based on polyvinyl alcohol or on cellulose.

Thus, those latices which have proved to be particularly advantageous for reducing the loss of mechanical properties after ageing are those based on a vinyl-type polymer with a protective colloid. Mention may be made, for example, of those based on a silanized or non-silanized vinyl chloride/olefin copolymer, especially a vinyl chloride/ethylene copolymer or more preferably a vinyl chloride/vinyl laurate/ethylene terpolymer.

As a variant, the dispersed phase may consist of the said polymer surrounded by surfactant, the surface-active molecule having a first end capable of being adsorbed on the surface of the polymer phase and a second end which is hydrophilic because of the suitable functional groups, the said functional groups coming together to form the surface of the dispersed particle. The surfactant may be chosen, in a manner known per se, from molecules capable of putting the said polymer in aqueous dispersion or emulsion.

Using the latices carrying hydrophilic groups that are mentioned above, the performance after ageing

of the insulation products according to the invention with regard to many mechanical properties, especially the puncture strength or the compressive strength, may be considerably improved.

5 However, introducing a very hydrophilic material into the insulation product may undesirably increase the amount of water capable of being held within the product, especially during storage in a wet atmosphere. In this case, it is possible to remedy this
10 drawback by adding a water-repellent agent to the latex, such as a silicone or fluorinated compounds, while still maintaining excellent mechanical properties after ageing.

15 A useful latex according to the method of the invention has advantageously a glass transition temperature T_g of less than 100°C , particularly of less than 80°C and especially of less than 50°C . Thus, it is thought possible that the polymer dispersed in the latex achieves sufficient plasticity in contact with
20 the filaments of mineral wool at the time of spraying the size and/or during passage through an oven to be fixed to the mineral wool in a manner compatible with the resin of the size. Since the minimum film-forming temperature T_m is generally less than the glass
25 transition temperature, it would also be possible for these latices to form a more or less uniform or continuous protective coating on the wool, which would preserve the material from attack by moisture.

30 Latices having a glass transition temperature T_g of greater than 80°C are generally not preferred since they form very rigid, or even brittle, deposits which are not beneficial for the desired mechanical properties.

35 Moreover, the glass transition temperature T_g of the latex is advantageously at least about -5°C , particularly at least 0°C and especially at least 5°C . Latices having a glass transition temperature T_g of less than -5°C form deposits which, in the dry state, are very soft, and even tacky, and which, because of

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the insulation products according to the invention with regard to many mechanical properties, especially the puncture strength or the compressive strength, may be considerably improved.

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35 of the latex is advantageously at least about -5°C , particularly at least 0°C and especially at least 5°C . Latices having a glass transition temperature T_g of less than -5°C form deposits which, in the dry state, are very soft, and even tacky, and which, because of

their lack of strength, do not have a significant effect on the mechanical properties after ageing in a wet medium.

The amount of latex added does not need to be very large in order to achieve a satisfactory level of improvement of the mechanical strength of the products. In general, the solids content of the latex introduced may advantageously be chosen to be less than 5% by weight with respect to the weight of the mineral wool, the latex being already effective at a content as low as 0.01%. In particular, the solids content of the latex introduced may be about 0.1 to 5%, especially 0.5 to 5%, by weight with respect to the weight of the mineral wool, but often a satisfactory result is obtained with a latex solids content of about 0.1 to 2%, or even about 0.1 to 1% and especially about 0.5 to 1%, with respect to the weight of the mineral wool.

With regard to the way of introducing the latex, the following alternative methods of implementation may be mentioned.

In a first method of implementation, the latex is mixed with the constituents of the size during the formulation of the latter, and then this modified sizing composition is applied in the usual manner to the mineral wool. This method of implementation is advantageous when the latex is compatible with the constituents of the size, i.e. the mixing does not cause any phase separation or any precipitation of solids.

In this regard, the subject of the invention is also a sizing composition for an insulation product, especially a thermal and/or acoustic insulation product, comprising a thermosetting resin and a latex.

The base formulation of the size may be as follows:

- thermosetting resin, particularly of the phenol-formaldehyde type:

50 to 90, especially 50 to 70,
parts by weight of solids;

- urea:

50 to 10, especially 50 to 30,
parts by weight:

5 with a resin + urea total = 100 parts by dry
weight;

- ammonium sulphate:

0 to 5, especially 1 to 3, parts by
weight;

- aqueous ammonia:

10 0 to 10, especially 2 to 10, parts
by weight (on the basis of NH_3);

- silane:

0 to 2 parts by weight;

- mineral oil:

15 0 to 20 parts by weight.

In a second method of implementation,
applicable even when the latex is not compatible, when
mixed, with the constituents of the size, the latex is
applied to the mineral wool separately from the size.
20 For example, the latex may be sprayed, as an aqueous
phase, close to the size-spraying ring, especially by
placing two superimposed spray rings in the path of the
mineral wool towards the shake-up device, one ring
(preferably the first ring in the direction of movement
25 of the wool) being intended for the latex and the other
ring being intended for the size.

Since the mineral wool thus sized is then
treated in an oven in order to cure the resin of the
size, care should be taken to ensure that the
30 conditions of the oven heat treatment (temperature and
residence time) do not affect the stability of the
latex. As a general rule, conventional manufacturing
conditions are completely compatible with the use of
latexes according to the invention.

35 The invention applies to insulation products
based on all types of mineral wool, both glass wool and
rock wool. It finds particularly useful application
when the product consists of a glass or rock wool
capable of dissolving in a physiological medium.

Examples of such materials are described for instance in EP-A-0,412,878, WO-A-95/31411, WO-A-95/32927, WO-A-93/22251, EP-A-0,459,897, WO-A-96/04213 and WO-A-95/31410.

5 These materials generally dissolve in a saline solution simulating a physiological medium at a rate of at least 30 and especially at least 40 or 50 ng/cm² per hour, measured at pH 4.5, and at a rate of at least 30 and especially at least 40 or 50 ng/cm² per hour,
10 measured at pH 7.5.

 Among the parameters which influence the water sensitivity of these materials, mention may be made of their relatively high content of alkali metal oxides, which may be about 8 to 25% and especially 14 to 20% by
15 weight in preferred glass compositions.

 To this should be added a boron oxide content of generally about 2 to 18%, particularly at least 4% or even at least 7% and especially from 4 to 13% or even from 7 to 15% by weight.

20 In particular, the sodium oxide Na₂O content may be greater than or equal to 16% by weight, for example about 16.5 to 19% by weight, with a potassium oxide content of about 0.2 to 0.5% by weight.

 In one particular example, according to
25 EP-A-412,878 the composition is as follows:

- SiO ₂	57 to 70%
- Al ₂ O ₃	0 to 5%
- CaO	5 to 10%
- MgO	0 to 5%
- Na ₂ O + K ₂ O	13 to 18%
- B ₂ O ₃	2 to 12%, especially 7 to 12%
- F	0 to 1.5%
- P ₂ O ₅	0 to 4%
- impurities	< 2%.

 In another particular example, according to WO-A-95/32927 the composition is as follows:

- SiO ₂	45 to 60%
- Al ₂ O ₃	< 2%
- CaO + MgO	10 to 16%
- Na ₂ O + K ₂ O	15 to 23%
- B ₂ O ₃	10 to 18%
- P ₂ O ₅	0 to 4%
- BaO	0 to 1%
- impurities	0 to 2%

5 In this regard, the subject of the invention is also an insulation product especially a thermal and/or acoustic insulation product, based on mineral wool provided with a size based on a thermosetting resin, especially a phenolic resin, in which the size contains a latex which improves the mechanical strength of the product after ageing, particularly in a wet medium, it being possible for this product to have any of the
10 above characteristics.

In general, an improved insulation product according to the invention may have the usual density characteristics, the density in general being at least 30 kg/m³. In a preferred embodiment, the insulation
15 product has a density of at least 50 kg/m³ and especially at least 80 kg/m³. These products, termed heavy products, are mainly used in roof-decking applications and are particularly exposed to moisture because of the thermal cycles and because of
20 atmospheric condensation. According to the invention, their mechanical strength after ageing under these conditions remains at a good level.

Finally, the subject of the invention is the use of a latex with a size for the insulation product, especially a thermal and/or acoustic insulation
25 product, based on mineral wool, in order to improve the mechanical strength after ageing, particularly in a wet medium, of the product.

Other features and advantages of the invention
30 will appear in the description of the detailed examples which follow.

EXAMPLE 1

Glass wool is manufactured using the internal centrifuging technique, in which the molten glass composition is converted into filaments by means of a tool called a centrifuging dish, comprising a basket forming the chamber for receiving the molten composition and a peripheral strip pierced with a multitude of holes; since the dish is rotated about its vertical axis of symmetry, the composition is thrown out through the holes under the action of the centrifugal force and the material escaping from the holes is drawn into filaments with the aid of a blast of drawing gas.

Conventionally, a size-spraying ring is placed underneath the fiberizing dishes so as to distribute the sizing composition uniformly over the glass wool which has just been formed.

The mineral wool thus sized is collected on a conveyor belt fitted with internal suction boxes which allow the mineral wool to be retained in the form of a felt or mat on the surface of the conveyor. The conveyor then travels through an oven where the resin of the size cures.

Compared with this conventional manufacturing technique, for the needs of this example, a second spray ring was mounted just above the sizing ring so as to spray, onto the wool, a latex composition which will be added to the size on the mineral filaments.

The composition of the glass (hereafter denoted G1) is of the type described in EP-A-0,412,878.

This is a so-called biosoluble glass, i.e. a glass capable of dissolving in a physiological medium. This type of glass is most particularly sensitive to exposure to liquid or atmospheric water for a prolonged period, it being possible for the hydrolytic attack of the glass to degrade the glass fibres with a potential loss of mechanical properties.

The composition (in parts by weight) of the size is as follows:

- phenol-formaldehyde resin 55 parts by weight of
R1 solids
(38% solids content by
weight, free phenol < 1.2%,
free formaldehyde < 7%):
- urea: 45 parts by weight
- mineral oil:
- ammonium sulphate: 3 parts by weight
- aqueous ammonia: 6 parts by weight
(on the basis of NH_3)
- silane: 0 to 1 parts by
weight.

5 The size is diluted with water before being
sprayed, the degree of dilution and the spray rate
being suitable for depositing about 7 to 15%, generally
about 7 to 10%, of dry matter with respect to the
weight of glass wool.

10 The latex sprayed above the size is, in the
case of this example, of the vinyl type. It is sold by
Wacker under the trade mark VINNOL and consists of an
aqueous dispersion of a vinyl chloride/vinyl
laurate/ethylene terpolymer stabilized by a protective
polyvinyl alcohol colloid. The polymer has a minimum
film-forming temperature of about 2°C and its glass
transition temperature is slightly higher. The aqueous
15 dispersion has a solids content of approximately 50%
and a pH of about 4.

20 A reference test was carried out without
spraying the latex, and two tests according to the
invention were carried out with an amount of latex
sprayed corresponding, respectively, to 1 and 2% of dry
matter with respect to the weight of glass. For all
these tests, as well as those of the following
examples, the amount of water provided by the latex is
taken into account in order to tailor the dilution of
25 the size in such a way that, with or without latex, the
glass wool receives the same amount of water.

The insulation product manufactured in this example is a panel having a density of about 80 kg/m^3 , the spraying of the latex in no way affecting achievement of the desired density.

5 The products obtained are subjected to measurements of the following characteristic parameters:

Immersion water uptake: This parameter characterizes the ability of the product to absorb water in the case of accidental contact with liquid water (during storage on site, for example). Indirectly, it makes it possible to quantify the degree of hydrophilicity of the additives of the glass wool.

15 It is measured by carrying out an immersion test according to a protocol similar to that in the ASTM C 240-72 standard.

20 A preweighed specimen is immersed, horizontally, in water at room temperature for 2 hours, with a water height of 30 mm above the level of the upper face of the specimen. Next, it is placed vertically on one edge in order to allow it to drain for a time of 15 minutes, after which it is weighed.

The results are expressed in terms of the mass of water in kg absorbed per m^3 of volume of material.

25 Friability: This parameter characterizes the loss of integrity of the mineral wool when the product is handled. In the context of the present invention, it is measured mainly in order to give an indication of the effectiveness of the latex deposit on the mineral filaments: a reduction in the friability is an indication that the latex has formed, at least on some of the filaments, a protective coating capable, should the filament break, of forming a kind of sheath retaining the glass in the insulation product.

35 It is measured in the following way: specimens are taken from the product in the form of cores 25 mm in diameter, these being cut up into approximately 3 mm slices. 3 grams of these samples are weighed out, the latter then being placed in a screen having a mesh size

of 1 mm together with 4 rubber balls. The screening operation is carried out with an amplitude of agitation of 2.5 mm for 3 minutes.

5 The material remaining in the screen is then weighed. The results are expressed as the percentage by weight of material which has passed through the meshes of the screen with respect to the initial weight.

10 The puncture strength (in N), the tear strength (in kPa) and the compressive strength (in kPa, for a 10% and 25% thickness reduction) are also measured in a conventional manner.

15 These properties are measured immediately after manufacture and after accelerated ageing (NORDTEST) for 7 days (168 hours) in an environmental chamber set to a temperature of 70°C and a relative humidity of 90 - 95%.

20 The results are given in Table 1 below, which shows, for each test, the percentage loss of property measured with respect to the initial state after manufacture and, for each test with a latex, the percentage improvement in the property in the aged state, calculated with respect to the reference without a latex, also in the aged state.

25 It is apparent from this that the latex used does not affect the water uptake capability of the product very much, showing a generally hydrophobic nature of the product. When the amount of latex reaches 2%, a limited hydrophilic tendency occurs, showing the relatively hydrophilic nature of the latex, due especially to the hydrophilic functional groups of the polyvinyl alcohol.

30 The latex seems to have been properly fixed to the glass wool, since the latter is made less friable than in the reference product.

35 It is surprising to note that the latex initially does not have a significant effect on the friability, but that a positive effect gradually appears with ageing of the product, resulting, after 7 days of ageing in the wet medium, in a much smaller

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loss than with the reference product. The integrity of the product according to the invention in the aged state is superior to the reference product by a factor of approximately two.

5

Table 1

Example 1 % latex/ glass	Immersion water uptake kg/m ³	Friability %			Puncture strength (N)				
		As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss		
0%, reference	19	2.7	20	17.3	178	61	-65%		
1% Vinnol	19	2.9	10 (-50%)	7.1	187	90 (+49%)	-52%		
2% Vinnol	23	3.5	11 (-45%)	7.5	184	90 (+49%)	-51%		
Example 1 % latex/ glass	Tear strength (kPa)			10% compressive strength (kPa)			25% compressive strength (kPa)		
	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss
0%, reference	14.2	3.4	-76%	19	12	-36%	50	23	-54%
1% Vinnol	13.6	5.5 (+62%)	-59%	20	13 (+8%)	-35%	47	27 (+17%)	-42%
2% Vinnol	11.4	4.8 (+41%)	-57%	19	13 (+8%)	-31%	45	27 (+17%)	-48%

With regard to the mechanical properties, it may be seen that the use of a latex has made it possible to reduce the loss of properties during ageing substantially, as well as to improve the level after ageing of each of the properties studied. The most remarkable improvements are in the puncture strength and tear strength.

15

Here again, it is surprising to note that the tear strength and the compressive strength (10% and 25%) are not improved, and are even slightly degraded, in the product immediately after its manufacture compared with the reference product. However, the incorporation of a latex makes the product much more stable over time in the sense that it suffers reductions in performance which are much less than in the reference product.

EXAMPLES 2 and 3

The above observations were confirmed by reproducing Example 1 with two other phenol-formaldehyde resins R2 and R3 having slightly different characteristics from resin R1.

In each case, a corresponding reference product without latex was prepared. The results obtained are given in Table 2 below.

Table 2

% latex/ glass	Immersion water uptake kg/m ³	Friability %			Puncture strength (N)		
		As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss
Example 2							
0%, reference	23	3.2	21	17.8	188	59	-69%
2% Vinnol	22	3.1	9.2 (-56%)	6.1	192	78 (+32%)	-59%
Example 3							
0%, reference	23	2.8	16.3	13.5	193	78	-59%
2% Vinnol	20	3.4	11.1 (-32%)	7.7	189	113 (+45%)	-40%

Table 2 (continued)

% latex/ glass	Tear strength (kPa)			10% compressive strength (kPa)			25% compressive strength (kPa)		
	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss
Example 2									
0%, reference	10.7	unmeas- urable	-100%	19	10	-47%	45	20	-55%
2% Vinnol	8.9	3	-66%	17	16 (+16%)	-6%	41	24 (+20%)	-56%
Example 3									
0%, reference	12.6	4.6	-63%	20	13	-36%	49	26	-47%
2% Vinnol	10.8	5.2 (+13%)	-52%	19	14 (+8%)	-31%	47	31 (+19%)	-34%

This again shows an improvement in all the properties of the same order of magnitude as in Example 1. It should be noted that Example 3 reaches a very high general level of performance.

EXAMPLES 4 and 5

These examples illustrate the repetition of Example 1 with other latices, respectively:

- Ex. 4: a vinyl acetate/methacrylic acid and ester copolymer having a minimum film-forming temperature of about 80°C (solids content of approximately 50%, pH of about 8);

- Ex. 5: a polyvinyl acetate homopolymer having a glass transition temperature of approximately 33°C (solids content of approximately 55%; pH of about 4 to 5).

These two latices are based on a polymer in which each monomer carries hydrophilic functional groups. As is apparent from Table 3 below, the polyvinyl acetate homopolymer is very hydrophilic, which means that the insulation product has a high water absorption. The supplementary addition of a silicone as a water-repellent agent, at a very low content of about 0.2%, makes it possible to bring the

absorption back to an acceptable level while maintaining the same improvement in the mechanical properties. The silicone may be substituted with a fluorinated-type water-repellent agent in order to reduce the water absorption while keeping the same advantageous level of mechanical properties.

Table 3

% latex/ glass	Immersion water uptake kg/m ³	Friability %			Puncture strength (N)				
		As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss		
Example 4									
0%, reference	23	2.8	21.5	18.7	192	46	-76%		
2% latex	67	2.5	18.7 (-13%)	16.2	199	65 (+41%)	-67%		
Example 5									
0%, reference	22	3.2	16.9	13.7	170	61	-64%		
2% latex	815	-	-	-	-	-	-		
2% latex + silicone	35	2.6	8.6 (-49%)	6	197	112 (+100%)	-43%		
% latex/ glass	Tear strength (kPa)			10% compressive strength (kPa)			25% compressive strength (kPa)		
	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss	As manu fact ured	Aged	Loss
Example 4									
0%, reference	16	5	-68%	18.6	15.3	-17%	50.1	25	-50%
2% latex	15.2	4.7 (+6%)	-69%	19.1	14.7 (-4%)	-23%	47	26.3 (+5%)	-44%
Example 5									
0%, reference	14	3.7	-81%	19	12	-36%	46	22	-52%
2% latex + silicone	13	7.7 (+108%	-40%	19	16 (+33%)	-16%	46	34 (+54%)	-26%

It may be seen that the hydrophilic latex of Example 4 has a very favourable effect on the puncture strength and tear strength, and has a lesser effect on the compressive strength. The very hydrophilic latex of Example 5 turns out to have an excellent capability of improving all the properties studied after ageing, although the behaviour just after manufacture is not improved.

10 EXAMPLE 6

Another insulation product is manufactured in the way indicated in Example 1 with the addition of 2% of latex, but with another glass (hereafter called G6) of the type described in WO-A-95/32927.

15 With this other glass, the addition of latex makes it possible to reduce the percentage loss of each of the properties studied. It will be noted in particular that there is a remarkable improvement in the puncture strength after ageing, which is actually
20 twice the strength of the reference without a latex.

EXAMPLE 7

Another insulation product is manufactured in the way indicated in Example 6, but with the polyvinyl acetate homopolymer latex used in Example 4, adding
25 1.5% of latex with respect to the weight of glass wool with 0.2% of silicone with respect to the weight of glass wool.

This hydrophilic latex makes it possible to achieve a more pronounced reduction in the percentage property loss than in Example 6. The level of each of
30 the mechanical properties is furthermore raised considerably, with in particular a 250% improvement in the puncture strength and a more than 90% improvement in the tear strength.

35 EXAMPLE 8

Another insulation product having a lower density of about 50 kg/m³ is manufactured under conditions which are otherwise identical to those in Example 2 (glass G1, resin R2, 2% of Vinnol latex).

In the case of this relatively light product, the mechanical strength is very markedly improved over the reference.

EXAMPLES 9 and 10

5 Other insulation products having a density of about 50 kg/m³ are manufactured with glass G6 under conditions which are similar to those in Example 6 and are adapted in order to modify the density of the product, the addition of Vinnol latex being only 1%.

10 In Example 9, resin R2 is used with resin/urea proportions in a ratio of 55/45 and the addition of latex is carried out in the upper spray.

The general level of mechanical strength is raised significantly over the reference product.

15 In Example 10, resin R1 is used, always with resin/urea proportions in a ratio of 55/45, the latex being mixed into the size, everything being applied to the glass wool using a single spray ring.

20 These new conditions of applying the latex are also conducive to improving the mechanical strength after ageing.

The results obtained for the products in Examples 6 to 10 are shown in Table 5 below.

EXAMPLES 11 and 12

25 As in Example 3, an insulation product having a density of approximately 80 kg/m³ is manufactured with resin R3, while adding only 1% of latex to the size by spraying with an upper ring and by using yet another glass (hereafter called G11) of the type described in WO-A-95/32927. Two new latices are used, namely:

- Ex 11: a silanized styrene/acrylic ester copolymer sold by Wacker under the reference Vinnapas LL6030 (film-forming temperature of 24°C);

35 - Ex 12: a vinyl chloride/ethylene copolymer sold by Wacker under the reference Vinnol CE 752 (film-forming temperature of 7°C). This is a hydrophobic latex.

40 These products are subjected to the same tests as those described above and the results are given in Table 4 below.

The invention, which has just been described in the particular case of insulation products based on glass wool of the type capable of dissolving in a physiological medium, is in no way limited to this embodiment. In particular, the latex may be used to improve the mechanical strength after ageing of products based on glasses that are less sensitive to exposure to water, or even products based on another mineral wool, such as rock wool. From the information given in the general description, those skilled in the art will be able to adapt the choice of materials depending on their particular requirements.

Table 4

	Immersion water uptake kg/m ³	Friability %			Puncture strength (N)				
		As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss		
0%, reference	18	1.2	16	14.8	277	41	-85%		
Example 11	16	1.5	10.9 (-32%)	9.4	221	57 (+39%)	-74%		
Example 12	11	1.3	11.1 (-30%)	9.4	229	47 (+15%)	-79%		
	Tear strength (kPa)			10% compressive strength (kPa)			25% compressive strength (kPa)		
	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss	As manu- fact- ured	Aged	Loss
0%, reference	21	7	-67%	26	18	-32%	55	23	-58%
Example 11	18	6.5 (-7%)	-63%	24	17 (-5%)	-30%	51	25 (+9%)	-51%
Example 12	20	6 (-14%)	-68%	24	19 (+5%)	-21%	55	27 (+17%)	-50%

Table 5

% latex/ glass	Friability %			Puncture strength (N)			Tear strength (kPa)			10% compressive strength (kPa)			25% compressive strengt (kPa)		
	As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss	As manufac- tured	Aged	Loss
Example 6															
0%, reference	4.7	20.5	15.8	127	36	-72%	15.1	6	-60%	22	17	-23%	47	28	-40%
2% latex	5.1	13.2	8.1	143	72	-50%	12	6.8	-43%	20	16.8	-16%	42.5	29	-31%
Example 7															
0%, reference	4.7	20.5	15.8	127	36	-72%	15.1	6	-60%	22	17	-23%	47	28	-40%
1.5% latex	2.6	5.4	2.8	176	128	-29%	13.8	11.6	-16%	25	21.7	-13%	50	39	-22%
Example 8															
0%, reference	2.2	8.6	6.4	90	30	-66%	8.4	4.7	-44%	12	7.1	-41%	23	11	-52%
2% latex	2.3	4.0	1.7	90	55	-39%	5.9	3.9	-34%	10.4	8	-18%	20	14	-30%
Example 9															
0%, reference	2.0	7.4	5.4	100	40	-60%	10.7	3.9	-63%	15.3	11	-28%	27	17	-37%
1% latex	2.0	4.7	2.7	95	55	-42%	10.3	7.9	-23%	15.1	14	-7%	28	21	-25%
Example 10															
0%, reference	1.3	8.1	6.8	100	55	-55%	7.5	2.5	-67%	11.4	8.4	-26%	23	13	-43%
1% latex	1.4	5.1	3.7	95	60	-36%	6.8	2.9	-57%	11.7	9.0	-23%	22	15	-32%

CLAIMS

1. Method of improving the mechanical strength after ageing, particularly in a wet medium, of an insulation product, especially a thermal and/or acoustic insulation product, based on mineral wool provided with a size comprising a thermosetting resin, especially a phenolic resin, in which method a latex is added to the size during the manufacture of the product.
2. Method according to Claim 1, in which the product is based on mineral wool capable of dissolving in a physiological medium, especially containing a proportion of alkali metal oxides of about 8 to 25% by weight of the wool.
3. Method according to Claim 1 or 2, in which the latex is based on an aqueous dispersion or emulsion of a polymer carrying hydrophilic functional groups, especially hydroxyl, carboxyl or ester functional groups.
4. Method according to Claim 3, in which the polymer is derived from one or more monomers each carrying at least one hydrophilic functional group.
5. Method according to Claim 3 or 4, in which the latex contains a polymer or copolymer which is of the vinyl type, especially a vinyl acetate homopolymer or copolymer, or of the acrylic type and/or which is derived from a carboxylic acid.
6. Method according to Claim 5, in which the latex is chosen from:
 - a polyvinyl acetate homopolymer, a vinyl acetate/(meth)acrylic acid or ester copolymer, a vinyl acetate/maleic ester copolymer, a vinyl acetate/olefin copolymer, a vinyl acetate/vinyl chloride copolymer;
 - a silanized acrylonitrile/acrylic ester or styrene/acrylic acid or ester copolymer.
7. Method according to one of Claims 1 to 3, in which the latex is based on an aqueous dispersion or emulsion of particles consisting of a polymer surrounded by a surfactant or by a protective colloid

having hydrophilic functional groups, especially one based on polyvinyl alcohol or on cellulose.

8. Method according to Claim 7, in which the latex comprises a silanized or non-silanized vinyl chloride/ethylene copolymer or a silanized or non-silanized vinyl chloride/vinyl laurate/ethylene terpolymer.

9. Method according to any one of Claims 3 to 8, in which a water-repellent agent, such as a silicone or a fluorinated compound, is added to the latex.

10. Method according to any one of the preceding claims, in which the latex is based on a polymer having a glass transition temperature T_g of less than 80°C and especially of less than 50°C .

11. Method according to any one of the preceding claims, in which the latex is based on a polymer having a glass transition temperature T_g of greater than -5°C and especially of greater than 0°C .

12. Method according to any one of the preceding claims, in which the solids content of the latex introduced is less than 5%, especially about 0.01 to 5%, by weight with respect to the weight of mineral wool.

13. Method according to any one of the preceding claims, in which the latex is mixed with the size before application to the mineral wool.

14. Method according to any one of Claims 1 to 12, in which the latex is applied to the mineral wool separately from the size.

15. Insulation product, especially a thermal and/or acoustic insulation product, based on mineral wool provided with a size based on a thermosetting resin, especially a phenolic resin, in which the size contains a latex which improves the mechanical strength of the product after ageing, particularly in a wet medium.

16. Insulation product according to Claim 15, in which the mineral wool consists of glass or rock wool capable of dissolving in a physiological medium,

especially containing a proportion of alkali metal oxides of about 8 to 25% by weight of the mineral wool.

17. Insulation product according to Claim 16, in which the mineral wool dissolves in a saline solution
- 5 simulating a physiological medium at a rate of at least 30 and especially at least 40 or 50 ng/cm² per hour, measured at pH 4.5, and at a rate of at least 30 and especially at least 40 or 50 ng/cm² per hour, measured at pH 7.5.
- 10 18. Insulation product according to one of Claims 15 to 17, which has a density of at least 30 kg/m³, especially at least 50 kg/m³ and particularly at least 80 kg/m³.
- 15 19. Use of a latex with a size for an insulation product, especially a thermal and/or acoustic insulation product, based on mineral wool, in order to improve the mechanical strength after ageing, particularly in a wet medium, of the product.
- 20 20. Use of a latex according to Claim 19, by mixing it in the size or by spraying it separately.
21. Sizing composition for an insulation product, especially a thermal and/or acoustic insulation product, comprising a thermosetting resin and a latex.

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PATENT**METHOD OF IMPROVING THE MECHANICAL STRENGTH OF AN
INSULATION PRODUCT BASED ON MINERAL WOOL, INSULATION
PRODUCT AND SIZING COMPOSITION**

Filed by: ISOVER SAINT-GOBAIN

ABSTRACT

In order to improve the mechanical strength after ageing, particularly in a wet medium, of an insulation product, especially a thermal and/or acoustic insulation product, based on mineral wool provided with a size, a latex is incorporated in the sizing composition which contains especially a phenolic resin.

The presence of the latex allows the loss of mechanical performance after ageing to be very considerably reduced.

The sizing composition advantageously comprises a latex based on an aqueous dispersion or emulsion of a polymer phase carrying hydrophilic functional groups.

Application to the manufacture of insulation products intended to be exposed to atmospheric condensation and/or based on mineral wool of lower hydrolytic resistance.

Fig. None

09/786113

Declaration and Power of Attorney for Patent Application

Déclaration et Pouvoirs pour Demande de Brevet

French Language Declaration

En tant l'inventeur nommé ci-après, je déclare par le présent acte que:

As a below named inventor, I hereby declare that:

Mon domicile, mon adresse postale et ma nationalité sont ceux figurant ci-dessous à côté de mon nom.

My residence, post office address and citizenship are as stated next to my name.

Je crois être le premier inventeur original et unique (si un seul nom est mentionné ci-dessous), ou l'un des premiers co-inventeurs originaux (si plusieurs noms sont mentionnés ci-dessous) de l'objet revendiqué, pour lequel une demande de brevet a été déposée concernant l'invention intitulée

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled

METHOD FOR MAKING A FIBROUS INSULATING

PRODUCT, SIZING STUFF AND COMPOSITION

(AS AMENDED)

et dont la description est fournie ci-joint à moins

the specification of which:

☐ ci-joint

☐ is attached hereto.

☐ a été déposée le _____

☒ was filed on August 30, 1999

sous le numéro de demande des Etats-Unis ou le numéro de demande international PCT

as United States Application Number or PCT International Application Number

_____ et modifiée le

PCT/FR99/02068 and was amended on

_____ (le cas échéant).

_____ (if applicable).

Je déclare par le présent acte avoir passé en revue et compris le contenu de la description ci-dessus, revendications comprises, telles que modifiées par toute modification dont il aura été fait référence ci-dessus.

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

Je reconnais devoir divulguer toute information pertinente à la brevetabilité, comme défini dans le Titre 37, § 1.56 du Code fédéral des réglementations.

I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, § 1.56.

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Je revendique par le présent acte avoir la priorité étrangère, en vertu du Titre 35, § 119(a)-(d) ou § 365(b) du Code des Etats-Unis, sur toute demande étrangère de brevet ou certificat d'inventeur ou, en vertu du Titre 35, § 365(a) du même Code, sur toute demande internationale PCT désignant au moins un pays autre que les Etats-Unis et figurant ci-dessous et, en cochant la case, j'ai aussi indiqué ci-dessous toute demande étrangère de brevet, tout certificat d'inventeur ou toute demande internationale PCT ayant une date de dépôt précédant celle de la demande à propos de laquelle une priorité est revendiquée.

Prior Foreign Application(s)

Demande(s) de brevet antérieure(s) dans un autre pays.

98/10923

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(Numéro)

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I hereby claim foreign priority under Title 35, United States Code, § 119(a)-(d) or § 365(b) of any foreign application(s) for patent or inventor's certificate, or § 365(a) of any PCT International application which designated at least one country other than the United States, listed below, and have also identified below, by checking the box, any foreign application for patent or inventor's certificate, or PCT International application having a filing date before that of the application on which priority is claimed.

Priority claimed
Droit de priorité
revendiqué

01 September 1998

(Day/Month/Year Filed)
(Jour/Mois/Année de dépôt)

☒

Yes
Oui

☐

No
Non

(Day/Month/Year Filed)
(Jour/Mois/Année de dépôt)

☐

Yes
Oui

☐

No
Non

Je revendique par le présent acte tout bénéfice, en vertu du Titre 35, § 119(e) du Code des Etats-Unis, de toute demande de Brevet provisoire effectuée aux Etats-Unis et figurant ci-dessous.

I hereby claim the benefit under Title 35, United States Code, § 119(e) of any United States provisional application(s) listed below.

(Application No.)
(N° de demande)

(Filing Date)
(Date de dépôt)

(Application No.)
(N° de demande)

(Filing Date)
(Date de dépôt)

Je revendique par le présent acte tout bénéfice, en vertu du Titre 35, § 120 du Code des Etats-Unis, de toute demande de brevet effectuée aux Etats-Unis, ou en vertu du Titre 35, § 365(c) du même Code, de toute demande internationale PCT désignant les Etats-Unis et figurant ci-dessous et, dans la mesure où l'objet de chacune des revendications de cette demande de brevet n'est pas divulgué dans la demande antérieure américaine ou internationale PCT, en vertu des dispositions du premier paragraphe du Titre 35, § 112 du Code des Etats-Unis, je reconnais devoir divulguer toute information pertinente à la brevetabilité, comme défini dans le Titre 37, § 1.56 du Code fédéral des réglementations, dont j'ai pu disposer entre la date de dépôt de la demande antérieure et la date de dépôt de la demande nationale ou internationale PCT de la présente demande.

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s), or § 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application.

PCT/FR99/02068

(Application No.)
(N° de demande)

30 August 1999

(Filing Date)
(Date de dépôt)

(Status) (patented, pending, abandoned)

(Statut) (breveté, en cours d'examen, abandonné)

(Application No.)
(N° de demande)

(Filing Date)
(Date de dépôt)

(Status) (patented, pending, abandoned)

(Statut) (breveté, en cours d'examen, abandonné)

Je déclare par le présent acte que toute déclaration ci-incluse est, à ma connaissance, véridique et que toute déclaration formulée à partir de renseignements ou de suppositions est tenue pour véridique; et de plus, que toutes ces déclarations ont été formulées en sachant que toute fausse déclaration volontaire ou son équivalent est passible d'une amende ou d'une incarcération, ou des deux, en vertu de la Section 1001 du Titre 18 du Code des Etats-Unis, et que de telles déclarations volontairement faussées risquent de compromettre la validité de la demande de brevet ou du brevet délivré à partir de celle-ci.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

French Language Declaration

POUVOIRS: En tant que l'inventeur cité, je désigne par la présente l'(les) avocat(s) et/ou agent(s) suivant(s) pour qu'ils poursuive(nt) la procédure de cette demande de brevet et traite(nt) toute affaire s'y rapportant avec l'Office des brevets et des marques: (mentionner le nom et le numéro d'enregistrement).

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith: (list name and registration number)



022850

Addresser toute correspondance à.

Send Correspondence to:



022850

Addresser tout appel téléphonique à:
(nom et numéro de téléphone)

Direct Telephone calls to: (name and telephone number)

(703) 413-3000

Nom complet de l'unique ou premier inventeur	Full name of sole or first inventor
1.00	Jacky JOACHIM
Signature de l'inventeur	Inventor's signature
Date	Date
	✓ <i>Joachim</i> 04/12/01
Domicile	Residence 14, rue de la Roche, F-60270
	Gouvieux, FRANCE
Nationalité	Citizenship
	France FRX
Adresse Postale	Post Office Address
	same as above
Nom complet du second co-inventeur, le cas échéant	Full name of second joint inventor, if any
2.00	Philippe ESPIARD
Signature de l'inventeur	Second inventor's signature
Date	Date
	<i>Espiard</i> April 12, 2001
Domicile	Residence 2, rue des Primeveres,
	F-60270 Gouvieux, FRANCE FRX
Nationalité	Citizenship
	France
Adresse Postale	Post Office Address
	same as above

(Fournir les mêmes renseignements et la signature de tout co-inventeur supplémentaire.)

(Supply similar information and signature for third and subsequent joint inventors.)

French Language Declaration

Nom complet du troisième co-inventeur, le cas échéant 3-00	Full name of third joint inventor, if any Bruno MAHEUXE
Signature de l'inventeur Date	Third inventor's signature Date ✓ 06/25/01
Domicile 16 rue des Templiers F-60290 NOUVELLY sous CLERMONT	Residence 16 rue du Donjon, F-60600 Clermont, FRANCE FRX
Nationalité	Citizenship France
Adresse Postale	Post Office Address same as above

Nom complet du quatrième co-inventeur, le cas échéant 4-00	Full name of fourth joint inventor, if any Roger GILBERT
Signature de l'inventeur Date	Fourth inventor's signature Date ✓ 04/02/01
Domicile	Residence 21, rue Anatole France, F-60290 Rantigny, FRANCE
Nationalité	Citizenship France FRX
Adresse Postale	Post Office Address same as above

Nom complet du cinquième co-inventeur, le cas échéant 5-00	Full name of fifth joint inventor, if any Wolfgang HOLSTEIN
Signature de l'inventeur Date	Fifth inventor's signature Date ✓ 05/09/01
Domicile	Residence Herdersgrasse 2, D-35315 Homburg, GERMANY
Nationalité	Citizenship Germany DEX
Adresse Postale	Post Office Address same as above

Nom complet du sixième co-inventeur, le cas échéant 6-00	Full name of sixth joint inventor, if any Gerald AMANT
Signature de l'inventeur Date	Sixth inventor's signature Date ✓ 06/13/01
Domicile	Residence Johann-Wilhelm Strasse 15, D-69259 Wilhelmsfeld, GERMANY
Nationalité	Citizenship Germany DEX
Adresse Postale	Post Office Address same as above

(Fournir les mêmes renseignements et la signature de tout co-inventeur supplémentaire.)

(Supply similar information and signature for third and subsequent joint inventors.)